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[Continued on next page]

(54) Title: THE PRODUCTION METHOD OF TRANSGENIC PORCINE PRODUCING HUMAN ERYTHROPOIETIN AND THE TRANSGENIC PORCINE

Preparation of Human Genomic EPO DNA



Construction of EPO Expression Vector

2.6 kb	2.5 kb	2.6 kb
Rat WAP promoter	hEPO genome	SV40 Poly A



DNA Microinjection



[a scene of microinjection]



[microinjected fertilized eggs]



Transplantation in Surrogate Mother Porcine and Parturition(Isolation of DNA from the Litters)



PCR Check



DNA Base Sequencing.

(57) Abstract: Disclosed are transgenic porcine capable of secreting human erythropoietin (EPO) in their milk and the preparation thereof. For the preparation of the transgenic porcine, a 2.6 kb WAP promoter from the mammary gland of a rat is first amplified by PCR. Along with this PCR product, the human EPO genome DNA fragment and an SV40 poly A DNA fragment are used to construct an expression vector. Separately, PMSG and human chorionic gonadotrophic (hCG) hormone are administered into porcine by intramuscular injection to induce porcine to ovulate excessively and the porcine were led to natural mating. From the porcine, the fertilized eggs in the first cell differentiation period are collected. Next, the expression vector is injected into male pronuclei which are immediately transplanted in surrogate mother porcine. The surrogate mother porcine are allowed to give birth to litters. Therefore, the present invention can produce the expensive medicine human EPO at low costs on a large scale, giving a contribution to the improving of human health.

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For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

THE PRODUCTION METHOD OF TRANSGENIC PORCINE PRODUCING HUMAN ERYTHROPOIETIN AND THE TRANSGENIC PORCINE

TECHNICAL FIELD

The present invention relates to transgenic porcine that are able to produce
5 human erythropoietin useful as a medicine. More particularly, the present
invention relates to transgenic porcine that are able to secrete human erythropoietin
in their milk, thereby producing the useful medicine at a low cost on a large scale
with stability. Also, the present invention is concerned with a method for
preparing such transgenic porcine.

10

BACKGROUND OF PRIOR ART

With an average life span of 120 days, human erythrocytes are generally destroyed at a level of one hundred-twentieth of their total number everyday in the reticuloendothelial system. However, they show homeostasis because they are newly produced equally every day (Guyton, Textbook of Medical Physiology,
15 pp56-60, W. B. Saunders Co., Philadelphia (1976)).

Erythrocytes are produced in the bone marrow through maturation and differentiation of erythroblasts during which the hormone EPO serves as a factor to stimulate the differentiation of less-differentiated cells into erythrocytes (Guyton, *supra*).

20

In the 1950s, EPO was found by observing the fact that a large amount of ^{59}Fe was incorporated into newly forming erythrocytes when sera of anemic animals were introduced into normal animals (Borsook, et al., Blood, 9, 734(1954)). A lack of oxygen or a shortage of erythrocytes owing to, for example, hemorrhage, or an increase of the number of anemic cells stimulates cells in the kidney of adults to synthesize and secrete increased amounts of erythropoietin into the bloodstream. This hormonal glycoprotein plays an important role in the control of erythropoiesis and the maintenance of the number

of erythrocytes in blood (Carnot et al., Compt. Rend. 143, 384 (1906); Kranz, S. B., Blood 77, 419(1991); Goldwasser, E., et al., in Peptide Growth Factors and their Receptors I, Sporn, M. B. and A. B. Roberts, eds., Springer-Verlag, Berlin, p. 747 (1990)).

5 As well known in the art, natural type EPO, which is responsible for the control of erythropoiesis, is secreted from the liver in fetuses. The secretion function for the EPO begins to move into the kidney at 120-140 days after the conception and the transferring of the secretion function is completed 40 days after the parturition. In adults, the kidney produces most of EPO while the liver is
10 responsible for the secretion of EPO at a level of 10% of the total amount secreted. In addition, a little amount of EPO is also known to be secreted in macrophages of the bone marrow.

15 EPO is maintained at a level of 15-30 mU per ml of blood or at a level of 0.01 mM in blood (Garcia, J. F., Lab. Clin. Med. 99, 624-635 (1982)). Higher levels of EPO in blood are measured from the patients suffering from aplastic anemia than from normal persons, so that the blood and/or urine of the patients are utilized to produce EPO (White, et al., Rec. Prog. Horm. Res. 16, 219 (196); Espada, et al., Biochem. Med. 3, 475 (1970); Fisher, pharmacol. Rev. 24, 459 (1972)).

20 As mentioned early, EPO is a glycoprotein with a molecular weight of about 30 kD, in which sugar chains are attached in N-glycosidic linkage to the 24th, the 38th and the 83rd amino acid residues and a sugar chain is attached in O-glycosidic linkage to the 126th amino acid residue (P. S. E. B. M. 216, 358-369 (1997)). Conventionally, EPO was produced in animal cells by a recombinant
25 technique, but at low amounts. In addition, the recombinant EPO suffers from the problems of being not identical in physiological functions to and of being poorer than natural type EPO.

30 EPO is very useful for the clinical treatment of anemic diseases, especially renal anemia and it is preferable that this therapeutic is prepared from human-derived materials owing to antigenicity. As mentioned early, EPO can be obtained by taking advantage of the blood or urine from patients suffering from

aplastic anemia. However, the amount of obtainable EPO from the patients, although being blood rich in EPO, is extremely limited.

From sera of sheep, EPO can be recovered in a stable water soluble form with a satisfactory titer, but this animal EPO includes the problem that it might act
5 as an antigen to the human body.

Biotechnology Co. Ltd., Cuba, took advantage of human erythropoietin (hEPO) cDNA to create a transgenic rabbit from which hEPO is secreted through its mammary gland. Likewise, Kuopioeogkr, Finland, was reported to have created a transgenic mouse capable of secreting hEPO via its mammary gland.
10 However, there have been found no reports which disclose transgenic porcine capable of secreting hEPO. Korean Pat. Publication No. 93-5917 describes that an hEPO gene is cloned and expressed in mammalian or insect cells. Not only is the EPO expressed only at a small amount in this process, but also glycosylation does not occur accurately so that the EPO is degraded rapidly in the body. In
15 Korean Pat. Appl'n No. 94-12082, an expression vector carrying a modified recombinant hEPO (rhEPO) is used to transform the animal cell COS-7 (ATCC CRL 1651, African green monkey kidney cell) into one which is able to produce rhEPO. This method, however, is unsuitable for large-scale production because of requiring continual transformation.

20 Korean Pat. No. 184778 discloses a method of producing rhEPO with stability and efficiency, which takes advantage of a permanent strain cell transfected by an expression vector carrying an hEPO gene. This patent is quite different from the present invention pertaining to the production of rhEPO in porcine milk.

25

DISCLOSURE OF THE INVENTION

Leading to the present invention, the intensive and thorough research on the production of human EPO, repeated by the present invention, resulted in the finding that a WAP promoter, in combination with SV40 Poly A, is very useful to incorporate a human EPO gene into the genomic DNA of porcine and the

recombinant expression vector can be used to create transgenic porcine which can secrete human EPO in their milk with stability.

Therefore, it is an object of the present invention to overcome the above problems encountered in prior arts and to provide transgenic porcine that are able 5 to secrete human EPO in their milk.

It is another object of the present invention to provide a method for preparing transgenic porcine capable of producing human EPO at low costs with stability.

In accordance with an embodiment of the present invention, there are 10 provided transgenic porcine (named "Saerome") capable of secreting human EPO in their milk with stability.

In accordance with another embodiment of the present invention, there is provided a method for preparing transgenic porcine capable of secreting human EPO in their milk, comprising the steps of: amplifying a 2.6 kb WAP promoter 15 from the mammary gland of a rat by a polymerase chain reaction; constructing an expression vector comprising a human erythropoietin genome DNA fragment and an SV40 poly A DNA fragment; administering PMSG and human chorionic gonadotrophic (hCG) hormone into porcine by intramuscular injection to induce porcine to ovulate excessively; determining the porcine as to their oestrus and 20 leading them to natural mating; collecting the fertilized eggs in the first cell differentiation period from the porcine; injecting the expression vector into male pronuclei and immediately transplanting them in surrogate mother porcine; allowing the surrogate mother porcine to give birth to litters; and identifying the incorporation of the base sequence of the Sequence List 1 into the genomic DNA 25 of the progeny.

BRIEF DESCRIPTION OF THE DRAWINGS

The above and other objects, features and other advantages of the present invention will be more clearly understood from the following detailed description taken in conjunction with the accompanying drawings, in which:

Fig. 1 is a schematic process flow showing the preparation of transgenic porcine which are able to secrete human EPO in their milk;

Fig. 2 shows the incorporation of human EPO gene into the genomic DNA of porcine through a polymerase chain reaction; and

5 Fig. 3 is a base sequence for a human EPO cDNA incorporated into the genomic DNA of porcine.

BEST MODES FOR CARRYING OUT THE INVENTION

A detail description will be given of a transgenic porcine capable of producing hEPO in its milk, below, in conjunction with the drawings. Before the present transgenic porcine capable of producing hEPO and preparation method thereof are disclosed or described, it is to be understood that explanation of well-known functions or structures might be eliminated if it is judged to make unclear the substance of the present invention. Also, it must be noted that the terminology used therein is defined with the purpose of describing particular embodiments only, but not limiting, and may be changed in its definition depending on the intention or usage of users. Therefore, it should be defined on the basis of the through-context of the present invention.

With reference to Fig. 1, there is schematically shown the entire procedure that allows the production of transgenic porcine capable of secreting hEPO in their milk. As a material to prepare a recombinant human EPO gene, we obtained a human genomic DNA fragment comprising an EPO gene from Prof. Kim. J. H., of the department of animal husbandry, Korean National KyoungSang University. Using a polymerase chain reaction (PCR), a 2.6 kb WAP promoter was amplified from a mammary gland gene of a rat, and the PCR product was cloned. Along 20 with an SV40 poly A gene and an hEPO gene, this promoter was used to construct a recombinant expression vector, which would serve as a DNA donor, as shown in 25 Table 1, below.

TABLE 1
EPO Expression Vector

DNAs	Rat WAP promoter	hEPO gene	SV40 Poly A
Size	2.6 kb	2.5 kb	2.6 kb

Porcine were allowed to ovulate excessively by the intramuscular injection of P.M.S.G (eCG) hormone, which is a superovulation-inducing hormone, and 5 human chorionic gonadotrophic (hCG) hormone. After the porcine were determined as to their oestrus and led to natural mating, the fertilized eggs in the first cell differentiation period were collected. The above expression vector was injected into male pronuclei which were immediately transplanted in surrogate mother porcine. One of the litters delivered from the surrogate mother porcine 10 was found to carry DNA fragments encoding human EPO as measured from its tail, blood and sperm by PCR. This result is given as shown in Fig. 2.

Given in the following Table 2 are the primer sequences which were used for the PCR for the determination as to whether the litters had the DNA fragments of interest.

15

TABLE 2

Primers	Sequences	Expected Sizes
Hepo-304	F 5'- CGA GAA TAT CAC GGT AGA ACC -3'	304 bp
	R 5'- CTC ATT CAA GCT GCA GTG TTC -3'	
Hepo-567	F 5'- AAG TGG TGC ATG GTG GTA GTC -3'	567 bp
	R 5'- TTA CAG AAA GGG CAA GCA GAA -3'	

Blood was taken from the EPO transgenic porcine and analyzed for erythrocyte properties. The results are given in Table 3, below.

TABLE 3

	No. of Erythrocytes (x10 ⁶ /ul)	Vol. Of Erythrocytes (%)
Control	4.63(100)	66.5(100)
Transformed	5.25(113)	78.3(118)

Electrophoresis of PCR products obtained from various copies of the genomic DNA of the litter delivered through the surrogate mother porcine gave information incorporated into the genomic DNA. Base sequencing analysis confirmed the incorporation, identifying the cDNA as having the base sequence shown in the following Base Sequence List. We named the resulting transgenic porcine "Saerome".

[SEQUENCE LIST]

Sequence No.: 1

10 Length of Sequence: 582

Type of Sequence: Nucleic Acids

Number of Strand: Double Strand

Topology: Linear

Type of Molecules: cDNA

15 Origin

EPO cDNA obtained from human liver DNA

Characteristics of Sequence

Mark representing a Characteristic: sig peptide

Position located: 1-81

20 Mark representing a Characteristic: mat peptide

Position located: 82-582

Mark representing a Characteristic: terminator

Position located: 580-582

[SEQUENCE 1]

	ATG CGG GTG CAC GAA TGT CCT GCC TGG CTG TGG CTT CTC CTG TCC	45
5	Met Gly Val His Glu Cys Pro Ala Trp Leu Trp Leu Leu Ser	
	-27	-20
	CTG CTG TCG CTC CCT CTG GGC CTC CCA GTC CTG GGC GCC CCA CCA	90
	Leu Leu Ser Leu Pro Leu Gly Leu Pro Val Leu Gly Ala Prp Pro	
	-10	+1
10	CGC CTC ATC TGT GAC AGC CGA GTC CTG GAG AGG TAC CTC TTG GAG	135
	Arg Leu Ile Cys Asp Ser Arg Val Leu Glu Arg Tyr Leu Leu Glu	
	10	
	GCC AAG GAG GCC GAG AAT ATC ACG ACG GGC TGT GCT GAA CAC TGC	180
	Ala Lys Glu Ala Glu Asn Ile Thr Thr Gly Cys Ala Glu His Cys	
15	20	30
	AGC TTG AAT GAG AAT ATC ACT GTC CCA GAC ACC AAA GTT AAT TTC	225
	Ser Leu Asn Glu Asn Ile Thr Val Pro Asp Thr Lys Val Asn Phe	
	40	
	TAT GCC TGG AAG AGG ATG GAG GTC GGG CAG CAG GCC GTA GAA GTC	270
20	Tyr Ala Trp Lys Arg Met Glu Val Gly Gln Gln Ala Val Glu Val	
	50	60
	TGG CAG GGC CTG GCC CTG TCG GAA GCT GTC CTG CGG GGC CAG	315
	Trp Gln Gly Leu Ala Leu Leu Ser Glu Ala Val Leu Arg Gly Gln	
	70	
25	GCC CTG TTG GTC AAC TCT TCC CAG CCG TGG GAG CCC CTG CAG CTG	360
	Ala Leu Leu Val Asn Ser Ser Gln Pro Trp Glu Pro Leu Gln Leu	
	80	90
	CAT GTG GAT AAA GCC GTC AGT GGC CTT CGC AGC CTC ACC ACT CTG	405
	His Val Asp Lys Ala Val Ser Gly Leu Arg Ser Leu Thr Thr Leu	
30	100	

CTT CGG GCT CTG GGA GCC CAG AAG GAA GCC ATC TCC CCT CCA GAT 450
Leu Arg Ala Leu Gly Ala Gln Lys Glu Ala Ile Ser Pro Pro Asp
110 120
GCG GCC TCA GCT GCT CCA CTC CGA ACA ATC ACT GCT GAC ACT TTC 495
5 Ala Ala Ser Ala Ala Pro Leu Arg Thr Ile Thr Ala Asp Thr Phe
130
CGC AAA CTC TTC CGA GTC TAC TCC AAT TTC CTC CGG GGA AAG CTG 540
Arg Lys Leu Phe Arg Val Tyr Ser Asn Phe Leu Arg Gly Lys Leu
140 150
10 AAG CTG TAC ACA GGG GAG GCC TGC AGG ACA GGG GAC AGA TGA 582
Lys Leu Tyr Thr Gly Gly Ala Cys Arg Thr Gly Asp Arg
160

As described hereinbefore, the present invention provides transgenic porcine capable of secreting human EPO in their milk, so that the expensive useful medicine can be produced at a low cost with stability on a large scale, thereby giving a contribution to the improving of human health.

The present invention has been described in an illustrative manner, and it is to be understood that the terminology used is intended to be in the nature of description rather than of limitation. Many modifications and variations of the present invention are possible in light of the above teachings. Therefore, it is to be understood that within the scope of the appended claims, the invention may be practiced otherwise than as specifically described.

CLAIMS

1. A method for preparing transgenic porcine capable of secreting human erythropoietin in their milk, comprising the steps of:

5 amplifying a 2.6 kb WAP promoter from the mammary gland of a rat by a polymerase chain reaction;

constructing an expression vector comprising a human erythropoietin genome DNA fragment, and an SV40 poly A DNA fragment;

administering PMSG and human chorionic gonadotrophic (hCG) hormone into porcine by intramuscular injection to induce porcine to ovulate excessively;

10 determining the porcine as to their oestrus and leading them to natural mating;

collecting the fertilized eggs in the first cell differentiation period from the porcine;

15 injecting the expression vector into male pronuclei and immediately transplanting them in surrogate mother porcine;

allowing the surrogate mother porcine to give birth to litters; and

identifying the incorporation of the base sequence of the Sequence List 1 into the genomic DNA of the progeny.

20 2. Transgenic porcine capable of producing human erythropoietin, prepared according to the method of claim 1.

3. The method as set forth in claim 1, wherein the expression vector comprises a 2.6 kb rat WAP promoter, a 2.5 kb hEPO and a 2.6 kb SV40 Poly A.

4. The method as set forth in claim 1, wherein the human erythropoietin cDNA comprises the base sequence shown in Fig. 3.

25 5. The transgenic porcine as set forth in claim 2, wherein the sperm DNA of the porcine comprises a gene coding for WAP-EPO.

6. The transgenic porcine as set forth in claim 2, wherein the human erythropoietin is WAP-EPO.

6'. The transgenic porcine as set forth in claim 2, wherein the human erythropoietin is produced in a form of WAP-EPO.

5 7. The transgenic porcine as set forth in claim 2, wherein the transgenic porcine is "Saerome".

8. The transgenic porcine as set forth in claim 2, wherein litters of the transgenic porcine have a WAP-EPO DNA.

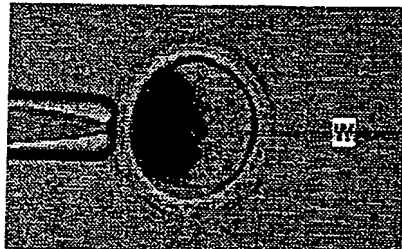
9. The transgenic porcine as set forth in any of claims 1 to 8, wherein the
10 produced erythropoietin can be readily used as a medicine.

1/3

[FIG. 1]

Preparation of Human Genomic EPO DNA**Construction of EPO Expression Vector**

2.6 kb	2.5 kb	2.6 kb
Rat WAP promoter	hEPO genome	SV40 Poly A

**DNA Microinjection**

[a scene of microinjection]

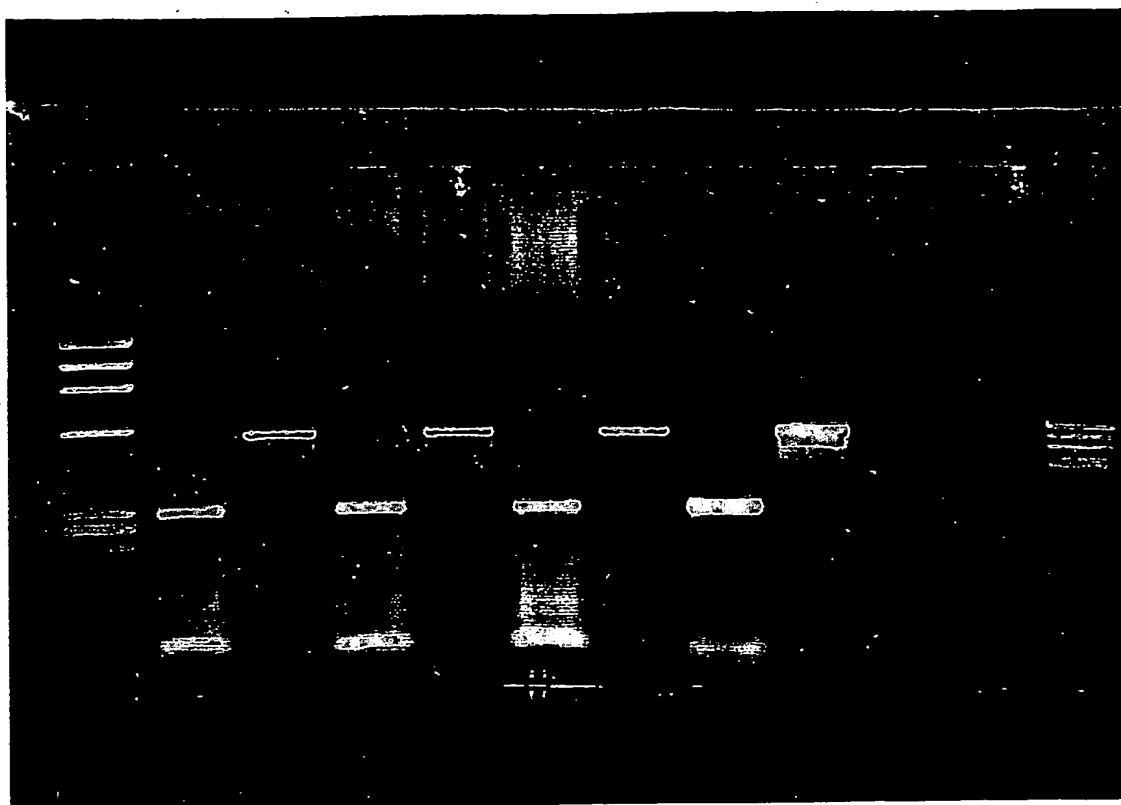


[microinjected fertilized eggs]

**Transplantation in Surrogate Mother Porcine and Parturition(Isolation of DNA from the Litters)****PCR Check****DNA Base Sequencing.**

2/3

[FIG. 2]



SM1

TAIL

BLOOD

SPERM

PC

NC

SM2

3 / 3

[FIG. 3]

ATG GGG GTG CAC GAA TGT CCT GCC TGG CTG TGG CTT CTC CTG TCC 45
 Met Gly Val His Glu Cys Pro Ala Trp Leu Trp Leu Leu Ser
 -27 -20
 CTG CTG TCG CTC CCT CTG GGC CTC CCA GTC CTG GGC GCC CCA CCA 90
 Leu Leu Ser Leu Pro Leu Gly Leu Pro Val Leu Gly Ala Prp Pro
 -10 +1
 CGC CTC ATC TGT GAC AGC CGA GTC CTG GAG AGG TAC CTC TTG GAG 135
 Arg Leu Ile Cys Asp Ser Arg Val Leu Glu Arg Tyr Leu Leu Glu
 10
 GCC AAG GAG GCC GAG AAT ATC ACG ACG GGC TGT GCT GAA CAC TGC 180
 Ala Lys Glu Ala Glu Asn Ile Thr Thr Gly Cys Ala Glu His Cys
 20 30
 AGC TTG AAT GAG AAT ATC ACT GTC CCA GAC ACC AAA GTT AAT TTC 225
 Ser Leu Asn Glu Asn Ile Thr Val Pro Asp Thr Lys Val Asn Phe
 40
 TAT GCC TGG AAG AGG ATG GAG GTC GGG CAG CAG GCC GTA GAA GTC 270
 Tyr Ala Trp Lys Arg Met Glu Val Gly Gln Gln Ala Val Glu Val
 50 60
 TGG CAG GGC CTG GCC CTG CTG TCG GAA GCT GTC CTG CGG GGC CAG 315
 Trp Gln Gly Leu Ala Leu Leu Ser Glu Ala Val Leu Arg Gly Gln
 70
 GCC CTG TTG GTC AAC TCT TCC CAG CCG TGG GAG CCC CTG CAG CTG 360
 Ala Leu Leu Val Asn Ser Ser Gln Pro Trp Glu Pro Leu Gln Leu
 80 90
 CAT GTG GAT AAA GCC GTC AGT GGC CTT CGC AGC CTC ACC ACT CTG 405
 His Val Asp Lys Ala Val Ser Gly Leu Arg Ser Leu Thr Thr Leu
 100
 CTT CGG GCT CTG GGA GCC CAG AAG GAA GCC ATC TCC OCT CCA GAT 450
 Leu Arg Ala Leu Gly Ala Gln Lys Glu Ala Ile Ser Pro Pro Asp
 110 120
 GCG GCC TCA GCT GCT CCA CTC CGA ACA ATC ACT GCT GAC ACT TTC 495
 Ala Ala Ser Ala Ala Pro Leu Arg Thr Ile Thr Ala Asp Thr Phe
 130
 CGC AAA CTC TTC CGA GTC TAC TCC AAT TTC CTC CGG GGA AAG CTG 540
 Arg Lys Leu Phe Arg Val Tyr Ser Asn Phe Leu Arg Gly Lys Leu
 140 150
 AAG CTG TAC ACA GGG GAG GCC TGC AGG ACA GGG GAC AGA TGA 582
 Lys Leu Tyr Thr Gly Gly Ala Cys Arg Thr Gly Asp Arg
 160

Sequence Listing

<110> Republic of Korea (Management:Rural Development Administration)

CHANG, Won-Kyong

PARK, Jin-Gi

SEONG, Hwan-Hoo

MIN, Kwan-Sik

YANG, Bo-Seok

IM, Gi-Sun

LEE, Yun-Keun

LEE, Chnag-Hyun

KIM, Jin-Hoei

<120> THE PROCUCTION METHOD OF TRANSGENIC PORCINE PRODUCING HUMAN ERYTHROPOIETIN AND THE TRANSGENIC PORCINE

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<141> 2000-02-14

<150> KR 10-2000-0006888

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<170> KopatentIn 1.55

<210> 1

<211> 582

<212> DNA

<213> TRANSGENIC PORCINE

<220>

<221> CDS

<222> (1)..(579)

<220>

<221> sig_peptide

<222> (1)..(81)

<223> Fix the number -1 for an amino acid of C end, give the decreasing number one by one to N end

<220>

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<222> (82)..(579)

<220>

<221> terminator

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ctg tcg ctc cct ctg ggc ctc cca gtc ctg ggc gcc cca cca cgc ctc 96
Leu Ser Leu Pro Leu Gly Leu Pro Val Leu Gly Ala Pro Pro Arg Leu
20 25 30

atc tgt gac agc cga gtc ctg gag agg tac ctc ttg gag gcc aag gag 144
Ile Cys Asp Ser Arg Val Leu Glu Arg Tyr Leu Leu Glu Ala Lys Glu
35 40 45

gcc gag aat atc acg acg ggc tgt gct gaa cac tgc agc ttg aat gag 192
Ala Glu Asn Ile Thr Thr Gly Cys Ala Glu His Cys Ser Leu Asn Glu
50 55 60

aat atc act gtc cca gac acc aaa gtt aat ttc tat gcc tgg aag agg 240
Asn Ile Thr Val Pro Asp Thr Lys Val Asn Phe Tyr Ala Trp Lys Arg
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85 90 95

ctg tcg gaa gct gtc ctg cgg ggc cag gcc ctg ttg gtc aac tct tcc 336
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100 105 110

cag ccg tgg gag ccc ctg cag ctg cat gtg gat aaa gcc gtc agt ggc 384
Gln Pro Trp Glu Pro Leu Gln Leu His Val Asp Lys Ala Val Ser Gly

115

120

125

ctt cgc agc ctc acc act ctg ctt cg^g gct ctg gga gcc cag aag gaa 432

Leu Arg Ser Leu Thr Thr Leu Leu Arg Ala Leu Gly Ala Gln Lys Glu

130

135

140

gcc atc tcc cct cca gat gc^g gcc tca gct gct cca ctc cga aca atc 480

Ala Ile Ser Pro Pro Asp Ala Ala Ser Ala Ala Pro Leu Arg Thr Ile

145

150

155

160

act gct gac act ttc cgc aaa ctc ttc cga gtc tac tcc aat ttc ctc 528

Thr Ala Asp Thr Phe Arg Lys Leu Phe Arg Val Tyr Ser Asn Phe Leu

165

170

175

cg^g gga aag ctg aag ctg tac aca ggg gag gcc tgc agg aca ggg gac 576

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185

190

aga t ga 582

Arg

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1

5

10

15

Leu Ser Leu Pro Leu Gly Leu Pro Val Leu Leu Gly Ala Pro Pro Arg Leu

20

25

30

Ile Cys Asp Ser Arg Val Leu Glu Arg Tyr Leu Leu Glu Ala Lys Glu

35

40

45

Ala Glu Asn Ile Thr Thr Gly Cys Ala Glu His Cys Ser Leu Asn Glu

50

55

60

Asn Ile Thr Val Pro Asp Thr Lys Val Asn Phe Tyr Ala Trp Lys Arg
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85 90 95

Leu Ser Glu Ala Val Leu Arg Gly Gln Ala Leu Leu Val Asn Ser Ser
100 105 110

Gln Pro Trp Glu Pro Leu Gln Leu His Val Asp Lys Ala Val Ser Gly
115 120 125

Leu Arg Ser Leu Thr Thr Leu Leu Arg Ala Leu Gly Ala Gln Lys Glu
130 135 140

Ala Ile Ser Pro Pro Asp Ala Ala Ser Ala Ala Pro Leu Arg Thr Ile
145 150 155 160

Thr Ala Asp Thr Phe Arg Lys Leu Phe Arg Val Tyr Ser Asn Phe Leu
165 170 175

Arg Gly Lys Leu Lys Leu Tyr Thr Gly Glu Ala Cys Arg Thr Gly Asp
180 185 190

Arg

INTERNATIONAL SEARCH REPORT

International application No.

PCT/KR00/00675

A. CLASSIFICATION OF SUBJECT MATTER**IPC7 C12N 5/06**

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

C12N 5/00

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
NCBI, pubmed, IBM patent database, USPTO patent database "Erythropoietin, transgenic"**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US 5955422 A (Kirin-Amgen, Inc) 21 Sept, 1999 (21. 09.199)	1-9
A	Proc. Natl. Acad. Sci. USA, 1990, 87:5178-5182.	1-9
A	Mol. Biol. Med., 1989, 5:255-261.	1-9
A	Transgenic Res, 1997, 6(1):75-84	1-9
A	DNA Cell Biol, 1999, 18(11):845-	1-9
A	Transgenic Res, 1998, 7 (4):311-7	1-9
A	Eur J. Biochem 1997, 245(2):482-9	1-9
A	Blood 1995, 85(10):2735-41	1-9
A	Biol Res 1995, 28(2):141-53	1-9

 Further documents are listed in the continuation of Box C. See patent family annex.

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Date of the actual completion of the international search

12 DECEMBER 2000 (12.12.2000)

Date of mailing of the international search report

13 DECEMBER 2000 (13.12.2000)

Name and mailing address of the ISA/KR

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